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IMPROVED METHOD FOR PREPARING FLOUR DOUGHS AND PRODUCTS MADE FROM SUCH DOUGHS USING GLYCEROL OXIDASE LOUGHS

FIELD OF THE INVENTION

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The present invention relates to the field of food manufacturing, in particular to the preparation of improved bakery products and other farinaceous food products. Specifically, the invention concerns the use of glycerol oxidase as a dough strengthening agent and improvement of the quality of baked and dried products made from such improved doughs. There is also provided a method of improving the properties of doughs and baked product by combined use of glycerol oxidase and a lipase.

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#### TECHNICAL BACKGROUND AND PRIOR ART

The "strength" or "weakness" of doughs is an important aspect of making farinaceous finished products from doughs, including baking. The "strength" or "weakness" of a dough is primarily determined by its content of protein and in particular the content and the quality of the gluten protein is an important factor in that respect. Flours with a low protein content is generally characterized as "weak". Thus, the

content is generally characterized as "weak". Thus, the cohesive, extensible, rubbery mass which is formed by mixing water and weak flour will usually be highly extensible when subjected to stress, but it will not return to its original dimensions when the stress is removed.

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Flours with a high protein content are generally characterized as "strong" flours and the mass formed by mixing such a flour and water will be less extensible than the mass formed from a weak flour, and stress which is applied during mixing will be restored without breakdown to a greater extent than is the case with a dough mass formed from a weak flour. Strong flour is generally preferred in most baking contexts because of the superior rheological and handling properties

2 of the dough and the superior form and texture qualities of the finished baked or dried products made from the strong flour dough. 5 Doughs made from strong flours are generally more stable. Stability of a dough is one of the most important characteristics of flour doughs. Within the bakery and milling industries it is known to use dough "conditioners" to strengthen the dough to increase its stability and strength. Such 10 dough conditioners are normally non-specific oxidizing agents such as e.g. iodates, peroxides, ascorbic acid, K-bromate or azodicarbonamide and they are added to dough with the aims of improving the baking performance of flour to achieve a dough with improved stretchability and thus having a desirable 15 strength and stability. The mechanism behind this effect of oxidizing agents is that the flour proteins, in particular gluten contains thiol groups which, when they become oxidized, form disulphide bonds whereby the protein forms a more stable matrix resulting in a better dough quality and impro-20 vements of the volume and crumb structure of the baked products. However, the use of several of the currently available nonspecific oxidizing agents is either objected to by consumers 25 or is not permitted by regulatory bodies. Hence it has been attempted to find alternatives to these conventional flour and dough additives, and the prior art has i.a. suggested the use of glucose oxidase and hexose oxidase for this purpose. 30 Glycerol oxidase is an oxidoreductase which is capable of oxidizing glycerol. Different types of glycerol oxidase have been described in the literature. Some of these glycerol oxidases need co-factors in order to oxidize glycerol (Shuen-Fu et al., 1996. Enzyme Micro. Technol., 18:383-387). 35 However, glycerol oxidase from Aspergillus japonicus does not require any co-factors in the oxidation of glycerol to glyceraldehyde (T. Uwajima and O. Terada, 1980. Agri. Biol.

3 Chem. 44:2039-2045). This glycerol oxidase has been characterized by T. Uwajima and O. Terada (Methods in Enzymology, 1982, 89:243-248) and 5 T. Uwajima et al. (Agric. Biol. Chem., 1979, 43:2633-2634), and has a pH optimum at 7.0 and  $K_{m}$  and  $V_{\text{max}}$  are 10.4 mM and 935.6  $\mu$ mol  $H_2O_2$  min<sup>-1</sup> respectively using glycerol as substrate. The enzyme is most active on glycerol but also other substrates like dihydroxyacetone, 1,3-propanediol, D-galac-10 tose ad D-fructose are oxidized by glycerol oxidase. Glycerol oxidase not requiring co-factors has also been isolated from Penicillium and characterized by Shuen-Fuh Lin et al. (Enzyme Micro. Technol., 1996, 18:383-387). This 15 enzyme has optimum activity in the pH range from 5.5 to 6.5 at 30°C. The enzyme is stable between 20 and 40°C but loses its activity at temperatures above 50°C. Other potential sources for glycerol oxidase according to the 20 invention include different fungal species as disclosed in DE-2817087-A, such as Aspergillus oryzae, Aspergillus parasiticus, Aspergillus flavus, Neurospora crassa, Neurospora sitophila, Neurospora tetrasperma, Penicillium nigricans, Penicillium funiculosum and Penicillium janthinellum. 25 Glycerol oxidase isolated from the above natural sources has been used for different applications. Thus, glycerol oxidase from Aspergillus japonicus has been used for glycoaldehyde production from ethylene glycol (Kimiyasu Isobe and Hiroshi 30 Nishise, 1995, Journal of Molecular Catalysis B: Enzymatic, 1:37-43). Glycerol oxidase has also been used in the combination with lipoprotein lipase for the determination of contaminated yolk in egg white (Yioshinori Mie, 1996. Food Research International, 29:81-84). DE-2817087-A and US 35 4,399,218 disclose the use of glycerol oxidase for the determination of glycerol.

4 It has now been found that the addition of a glycerol oxidase to a flour dough results in an increased resistance hereof to deformation when the dough is stretched, i.e. this enzyme confers to the dough an increased strength whereby the dough 5 becomes less prone to mechanical deformation. Accordingly, glycerol oxidase is highly useful as a dough conditioning agent in the manufacturing of flour dough based products including not only bread products but also other products made from flour doughs such as noodles and alimentary paste 10 products. It has also been found that the dough strengthening effect of glycerol oxidase is potentiated significantly when it is combined with a lipase, which in itself does not affect the 15 dough strength. Furthermore, the combined use of glycerol oxidase and lipase results in an improvement of bread quality, in particular in respect of specific volume and crumb homogeneity, which is not a simple additive effect, but reflects a synergistic effect of these two types of enzymes. 20 SUMMARY OF THE INVENTION Accordingly, the invention relates in a first aspect to a 25 method of improving the rheological properties of a flour dough and the quality of the finished product made from the dough, comprising adding to the dough 10 to 10,000 units of a glycerol oxidase per kg of flour. 30 In a further aspect there is provided a method of improving the rheological properties of a flour dough and the quality of the finished product made from the dough, comprising adding to the dough a glycerol oxidase and a lipase. 35 The invention pertains in a still further aspect to dough improving composition comprising a glycerol oxidase and at least one further dough ingredient or dough additive.

15 The expression "rheological properties" as used herein refers particularly to the effects of dough conditioners on dough strength and stability as the most important characteristics of flour doughs. According to American Association of Cereal Chemists (AACC) Method 36-01A the term "stability" can be

20 defined as "the range of dough time over which a positive response is obtained and that property of a rounded dough by which it resists flattening under its own weight over a course of time". According to the same method, the term "response" is defined as "the reaction of dough to a known 25 and specific stimulus, substance or set of conditions, usually determined by baking it in comparison with a control"

As it is mentioned above, it is generally desirable to improve the baking performance of flour to achieve a dough with improved stretchability and thus having a desirable strength and stability by adding oxidizing agents which cause the formation of protein disulphide bonds whereby the protein forms a more stable matrix resulting in a better dough quality and improvements of the volume and crumb structure of baked products.

Thus, the term "rheological properties" relates to the above physical and chemical phenomena which in combination will

7 nally used dough components such as salt, sweetening agents such as sugars, syrups or artificial sweetening agents, lipid substances including shortening, margarine, butter or an animal or vegetable oil, glycerol and one or more dough 5 additives such as emulsifying agents, starch degrading enzymes, cellulose or hemicellulose degrading enzymes, proteases, lipases, non-specific oxidizing agents such as those mentioned above, flavouring agents, lactic acid bacterial cultures, vitamins, minerals, hydrocolloids such as algina-10 tes, carrageenans, pectins, vegetable gums including e.g. guar gum and locust bean gum, and dietary fiber substances. Conventional emulsifying agents used in making flour dough products include as examples monoglycerides, diacetyl tar-15 taric acid esters of mono- and diglycerides of fatty acids, and lecithins e.g. obtained from soya. Among starch degrading enzymes, amylases are particularly useful as dough improving additives. Other useful starch degrading enzymes which may be added to a dough composition include glucoamylases and pullu-20 lanases. In the present context, further interesting enzymes are xylanases and oxidoreductases such as glucose oxidase, pyranose oxidase, hexose oxidase, sulfhydryl oxidase, and lipases. 25 A preferred flour is wheat flour, but doughs comprising flour derived from other cereal species such as from rice, maize, barley, rye and durra are also contemplated. In accordance with the invention, the dough is prepared by 30 admixing flour, water, the glycerol oxidase and optionally other ingredients and additives. The glycerol oxidase can be added together with any dough ingredient including the water or dough ingredient mixture or with any additive or additive mixture. The dough can be prepared by any conventional dough 35 preparation method common in the baking industry or in any other industry making flour dough based products.

8 The glycerol oxidase can be added as a liquid preparation or in the form of a dry powder composition either comprising the enzyme as the sole active component or in admixture with one or more other dough ingredients or additive. 5 The amount of the glycerol oxidase added is an amount which results in the presence in the dough of 10 to 5,000 units (as defined in the following) such as 10 to 2,500 units per kg of flour. In useful embodiments, the amount is in the range of 10 20 to 1,500 units per kg of flour. The effect of the glycerol oxidase on the rheological properties of the dough can be measured by standard methods according to the International Association of Cereal Chemi-15 stry (ICC) and the American Association of Cereal Chemistry (AACC) including the amylograph method (ICC 126), the farinograph method (AACC 54-21) and the extensigraph method (AACC 54-10). The AACC method 54-10 defines the extensigraph in the following manner: "the extensigraph records a load-extension 20 curve for a test piece of dough until it breaks. Characteristics of load-extension curves or extensigrams are used to assess general quality of flour and its responses to improving agents". In effect, the extensigraph method measures the relative strength of a dough. A strong dough exhibits a 25 higher and, in some cases, a longer extensigraph curve than does a weak dough. In a preferred embodiment of the method according to the invention, the resistance to extension of the dough in terms 30 of the ratio between the resistance to extension (height of curve, B) and the extensibility (length of curve, C), i.e. the B/C ratio as measured by the AACC method 54-10 is increased by at least 10% relative to that of an otherwise similar dough not containing glycerol oxidase. In more pre-35 ferred embodiments, the resistance to extension is increased by at least 20%, such as at least 50% and in particular by at least 100%.

11 tion with glycerol, will have a significant improving effect on the extensibility and stability hereof resulting in finished paste product having improved textural and eating qualities. In a useful embodiment, there is provided a dough improving method wherein at least one further enzyme is added to the dough ingredient, dough additive or the dough. In the present context, suitable enzymes include cellulases, hemicellulases, 10 xylanases, starch degrading enzymes, oxidoreductases and proteases. In a further aspect, the invention relates to a method of improving the rheological properties of a flour dough and the 15 quality of the finished products made from the dough which comprises that both a glycerol oxidase and a lipase is added to the dough. It was surprisingly found that the two types of enzymes were 20 capable of interacting with each other under the dough conditions to an extent where the effect on improvement of the dough strength and bread quality by the enzymes was not only additive, but the effect was synergistic. 25 Thus, with respect to improvement of dough strength it was found that with glycerol oxidase alone, the B/C ratio as measured after 45 minutes of resting was increased by 34%, with lipase alone no effect was observed. However, when combining the two enzymes, the B/C ratio was increased by 30 54%, i.e. combining the glycerol oxidase with the lipase enhanced the dough strengthening effect of glycerol oxidase by more than 50%. Thus, one objective of combining glycerol

oxidase and a lipase is to provide an enhancement of the dough strengthening effect of glycerol oxidase by at least

35 25% such as at least 50% including at least 75%, determined as described herein.

In relation to improvement of finished product, it was found

that the combined addition of glycerol oxidase and a lipase resulted in a substantial synergistic effect in respect to crumb homogeneity as defined herein. Also, with respect to the specific volume of baked product a synergistic effect was found. Thus, for a bread product, the addition of lipase alone typically results in a negligible increase of the specific volume, addition of glycerol oxidase alone in an increase of about 25%, whereas a combined addition of the two enzymes results in an increase of more than 30%.

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Further in relation to improvement of the finished product, it was found that the addition of lipase resulted in modification of the glycolipids, monogalactosyl diglyceride and digalactosyl diglyceride present in dough. These components were converted to the more polar components monogalactosyl monoglyceride and digalactosyl monoglyceride. As galactosyl monoglycerides are more surface active components than galactosyl diglycerides it is assumed that galactosyl monoglycerides contributed to the observed improved crumb cell structure and homogeneity. Thus, one objective of using lipase is to hydolyse at least 10% of the galactosyl diglycerides normally present in a flour dough to the corresponding galactosyl monoglycerides, such as at least 50% including at least 100%.

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The details of such a method using combined addition of glycerol oxidase and lipase are, apart from the use of a lipase in combination with glycerol oxidase, substantially similar to those described above for a method according to the invention which does not require the addition of a lipase.

When using, in accordance with the invention, a lipase in combination with a glycerol oxidase, the amount of lipase is 35 typically in the range of 10 to 100,000 lipase units (LUS) (as defined in the following) per kg flour including the range of 10 to 20,000 LUS, e.g. 100 to 15,000 LUS such as 500 to 10,000 LUS.

Lipases that are useful in the present invention can be derived from a bacterial species, a fungal species, a yeast species, an animal cell and a plant cell. Whereas the enzyme 5 may be provided by cultivating cultures of such source organisms naturally producing lipase, it may be more convenient and cost-effective to produce it by means of genetically modified cells such as it is described in details in the following examples. In the latter case, the term "derived" 10 may imply that a gene coding for the lipase is isolated from a source organism and inserted into a host cell capable of expressing the gene.

Thus, the enzyme may in a useful embodiment be derived from 15 an Aspergillus species including as examples A. tubigensis, A. oryzae and A. niger.

Presently preferred lipases include the lipase designated Lipase 3, the production and characteristics of which is 20 described in details in the following examples, or a mutant of this enzyme. In the present context, the term "mutant" refers to a lipase having, relative to the wild-type enzyme, an altered amino acid sequence. A further preferred lipase is the lipase found in the commercial product, GRINDAMYLTM EXEL 25 16.

In a further aspect of the invention there is provided a dough improving composition comprising a glycerol oxidase and at least one further dough ingredient or dough additive.

The further ingredient or additive can be any of the ingredients or additives which are described above. The composition may conveniently be a liquid preparation comprising the glycerol oxidase. However, the composition is conveniently in 35 the form of a dry composition.

The amount of the glycerol oxidase in the composition is in the range of 10 to 10,000 units per kg flour. It will be

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BRIEF DESCRIPTION OF THE FIGURES

5 The present invention is further illustrated by reference to the accompanying figures in which

Fig. 1 shows the restriction map of the genomic clone of the lipA gene,

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Fig. 2 shows the structure of the lipA gene encoding lipase 3,

Fig. 3 shows a chromatogram of HIC fractionated culture 15 supernatant of an Aspergillus tubigensis transformant with 62-fold increase of lipase 3, and

Fig. 4 shows a chromatogram of HIC fractionated culture supernatant of the untransformed *Aspergillus tubigensis* 20 strain.

The invention will now be described by way of illustration in the following non-limiting examples.

25 A. PRODUCTION AND PURIFICATION OF GLYCEROL OXIDASE (GLOX)

EXAMPLE 1

Production, extraction and purification of glycerol oxidase
30 using different strains and cultivation conditions

1. Production, extraction and purification of glycerol oxidase using Aspergillus japonicus ATCC 1042 cultivated in a production medium containing 3% glycerol

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The following assay for determination of glycerol oxidase activity was used:

The assay is based on the method described by Sullivan and Ikawa (Biochimica and Biophysica Acta, 1973, 309:11-22), by modified as described in the following. An assay mixture containing 150 µl 2% glycerol (in 100 mM phosphate buffer,

Ikawa (Biochimica and Biophysica Acta, 1973, 309:11-22), but modified as described in the following. An assay mixture containing 150 μl 2% glycerol (in 100 mM phosphate buffer, pH 5 7.0), 120 μl 100 mM phosphate buffer, pH 7.0, 10 μl o-dianisidin dihydrochloride (Sigma D 3252, 3 mg/ml in H<sub>2</sub>O), 10 μl peroxidase (POD) (Sigma P8125, 0.1 mg/ml in 100 mM phosphate buffer, pH 7.0) and 10 μl glycerol oxidase (GLOX) solution. The controls are made by adding buffer in place of 10 GLOX solution. The incubation is started by the addition of glycerol. After 15 minutes of incubation at 25°C in microtiter plates, the absorbance at 402 nm is read in a Elisa reader. A standard curve is constructed using varying concentrations of H<sub>2</sub>O<sub>2</sub> in place of the enzyme solution. The rea-

15 ction can be described in the following manner:

GLOX

Glycerol +  $O_2$  ----> glyceraldehyde +  $H_2O_2$ 

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POD

 $H_2O_2$  + o-dianisidine<sub>red</sub> -----> 2  $H_2O$  + o-dianisidine<sub>ox</sub>

Oxidised o-dianisidine has a yellow colour absorbing at 402 25 nm.

One glycerol oxidase unit (U) is the amount of enzyme which catalyses the production of 1  $\mu$ mole  $H_2O_2$  per minute at 25°C, pH 7.0 at a substrate concentration of 0.2 M glycerol.

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A spore suspension of Aspergillus japonicus ATCC 1042 was prepared by incubating A.japonicus on PDA medium (30°C, 7 days) and washing with 10 ml of 0.2% Tween 80. A preculture was prepared by inoculating 1 ml of the resulting spore 35 suspension in 300 ml production medium containing 3.0 % of glycerol (87%, Merck), 0.3% of yeast extract (Difco), 0.1% of meat extract (Difco), 0.1 % KH<sub>2</sub>PO<sub>4</sub> (Merck), 0.1% of MGSO<sub>4</sub> \*

2. Production, extraction and purification of glycerol oxidase using Aspergillus japonicus ATCC 1042 cultivated in a production medium containing 5% glycerol

30 supernatant contained 8.7 units of glycerol oxidase activity

per ml.

A spore suspension of Aspergillus japonicus ATCC 1042 was

prepared as described above. A preculture was prepared by inoculating 1 ml of the resulting spore suspension into a flask (500 ml) containing 200 ml production medium (5.0 % glycerol, 0.25% yeast extract, 0.1% Malt extract, 0.7% anti-

- 5 foam (Contra spum), pH adjusted to 6.2 with HCl, sterilization at 121°C for 90 minutes). The preculture was incubated 3 days at 30°C with continuous shaking (200 rpm).
  - A 6 litre fermenter with 5 litre production medium as described above was inoculated with 50 ml of the resulting
- 10 preculture and cultured at 30°C for 3 days under continuous stirring (250 rpm) and aeration (5 l/min). After culturing the mycelia was harvested from the resulting culture broth by filtration on a Whatman GF/B filter by suction, and washed with 3 litre ionized water containing 0.9% NaCl.
- 15 The resulting mycelia mat was frozen in liquid nitrogen, suspended in 200 ml of 50 mM phosphate buffer (pH 7.0) and disrupted by ultrasonication (Branson, Sonifer 250) at 5°C (4 minutes). After disruption, the mycelia was removed by filtration on a Whatman GF/A filter by suction. The enzyme in
- 20 the resulting filtrate was concentrated on a AMICON 8400 ultrafiltration unit and contained 87 units of glycerol oxidase per ml after ultrafiltration.
- 3. Production, extraction and purification of glycerol

  oxidase using Aspergillus japonicus ATCC 1042 cultivated in a
  production medium containing 10% glycerol

A spore suspension of Aspergillus japonicus ATCC 1042 was prepared as described above. A 1 ml sample of the resulting 30 spore suspension was inoculated into each of 5 flasks (500 ml) with 200 ml production medium containing 10.0 % of glycerol, 0.1% of yeast extract and 0.1% of malt extract (pH adjusted to 6.2 with HCl, sterilization at 121°C for 15 minutes). The cultures were incubated for 5 days at 30°C with 35 shaking (140 rpm).

The extraction and concentration of the enzyme was carried

20 out as described above. The resulting filtrate contained 66 units of glycerol oxidase per ml after ultrafiltration. 4. Production of glycerol oxidase from Penicillium funi-5 culosum and Penicillium janthinellum Spore suspensions of Penicillium funiculosum NRRL 1132 and Penicillium janthinellum NRRL 2016 were prepared as described above. A 1 ml sample of each of the resulting spore suspen-10 sions was inoculated into separate flasks (1000 ml) containing 100 g wheat bran and 100 ml water (two flasks for each culture) Glycerol oxidase was extracted by suspending the wheat bran 15 cultures in 900 ml of 30 mM phosphate buffer (pH 6.5) containing 0.1% Triton X100 (Merck). The mycelial mat was removed from the cultivation media by filtration using a Whatman GF/B filter. The resulting mycelia mat was frozen in liquid nitrogen, suspended in 200 ml of 50 mM phosphate 20 buffer (pH 7.0) and disrupted by ultrasonication (Branson, Sonifer 250) at 5°C (4 minutes). After disruption, the mycelia was removed by filtration on a Whatman GF/A filter by suction. The resulting filtrate from the Penicillium funiculosum culture contained 7.4 units of glycerol oxidase per ml, 25 and the resulting filtrate from the Penicillium janthinellum culture contained 11.3 units of glycerol oxidase per ml. B.PRODUCTION, PURIFICATION AND CHARACTERIZATION OF ASPERGILLUS TUBIGENSIS LIPASE 3 30 Materials and Methods (i) Determination of lipase activity and protein 35 1. Plate assay on tributyrin-containing medium

The assay is modified from Kouker and Jaeger (Appl. Environ. Microbiol., 1987, 53:211-213).

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A typical protocol for this assay is as follows: 100 ml 2% agar in 50 mM sodium phosphate buffer (pH 6.3) is heated to boiling, and after cooling to about 70°C under stirring, 5 ml 5 0.2% Rhodamine B is added under stirring plus 40 ml of tributyrin. The stirring is continued for 2 minutes. The mixture is then sonicated for 1 minute. After an additional 2 minutes of stirring, 20 ml of the agar mixture is poured into individual petri dishes. In the absence of lipase activity, the 10 agar plates containing tributyrin and Rhodamine B will appear opaque and are pink coloured.

To quantify lipase activity, holes having a diameter of 3 mm are punched in the above agar and filled with 10 µl of lipase 15 preparation. The plates are incubated for varying times at 37°C. When lipase activity is present in the applied preparation to be tested, a sharp pink/reddish zone is formed around the holes. When the plates are irradiated with UV light at 350 nm, the lipase activity is observed as halos of orange 20 coloured fluorescence.

#### 2. Modified Food Chemical Codex assay for lipase activity

Lipase activity based on hydrolysis of tributyrin is measured according to Food Chemical Codex, Forth Edition, National Academy Press, 1996, p. 803. With the modification that the pH is 5.5 instead of 7. One LUT (lipase unit tributyrin) is defined as the amount of enzyme which can release 2  $\mu$ mol butyric acid per min. under the above assay conditions.

#### 3. p-nitrophenyl acetate assay

Lipase activity can also be determined colorimetrically using p-nitrophenyl acetate as a substrate e.g. using the following 35 protocol: In a microtiter plate 10  $\mu$ l of sample or blank is added followed by the addition of 250  $\mu$ l substrate (0.5 mg p-nitrophenyl acetate per ml 50 mM phosphate buffer, pH 6.0).

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Lipase activity can be determined by using p-nitrophenyl hexanoate as a substrate. This assay is carried out by adding 10 10 µl of sample preparation or blank to a microtiter plate followed by the addition of 250 µl substrate (0.5 mg p-nitrophenyl hexanoate per ml of 20 mM phosphate buffer, pH 6.). At this concentration of substrate the reaction mixture appears as a milky solution. The microtiter plate is incubated for 5 minutes at 30°C and the absorbance at 405 nm is read in a microplate reader.

#### 5. Titrimetric assay of lipase activity

20 Alternatively, lipase activity is determined according to Food Chemical Codex (3rd Ed., 1981, pp 492-493) modified to sunflower oil and pH 5.5 instead of olive oil and pH 6.5. The lipase activity is measured as LUS (lipase units sunflower) where 1 LUS is defined as the quantity of enzyme 25 which can release 1  $\mu$ mol of fatty acids per minute from sunflower oil under the above assay conditions.

#### 6. Protein measurement

30 During the course of purification of lipase as described in the following, the protein eluted from the columns was measured by determining absorbance at 280 nm. The protein in the pooled samples was determined in microtiter plates by a sensitive Bradford method according to Bio-Rad (Bio-Rad 35 Bulletin 1177 EG, 1984). Bovine serum albumin was used as a standard.

#### EXAMPLE 2

## Production, purification and characterization of lipase 3

#### 5 2.1. Production

A mutant strain of Aspergillus tubigensis was selected and used for the production of wild type lipase. This lipase is referred to herein as lipase 3. The strain was subjected to a

- 10 fermentation in a 750 l fermenter containing 410.0 kg of tap water, 10.8 kg soy flour, 11.1 kg ammonium monohydrogenphosphate, 4.0 kg phosphoric acid (75%), 2.7 kg magnesium sulfate, 10.8 kg sunflower oil and 1.7 kg antifoam 1510. The substrate was heat treated at 121°C for 45 minutes. The
- 15 culture media was inoculated directly with  $7.5\times10^9$  spores of the mutant strain. The strain was cultivated for three days at 38°C, pH controlled at 6.5, aeration at 290 l/min and stirring at 180 rpm the first two days and at 360 rpm the last day. The fermentate was separated using a drum filter
- 20 and the culture filtrate was concentrated 3.8 times by ultrafiltration. The concentrated filtrate was preserved with potassium sorbate (0.1%) and sodium benzoate (0.2%) and used as a starting material for purification of lipase.

### 25 2.2. Purification of lipase

A 60 ml sample of ferment (cf. 2.1) containing 557 LUS/ml, pH 5.5 was first filtered through a GF/B filter and subsequently through a 0.45  $\mu$ m filter. The filtered sample was desalted

- 30 using a Superdex G25 SP column (430 ml, 22 x 5 cm) equilibrated in 20 mM triethanolamine, pH 7.3. The flow rate was 5 ml/min. The total volume after desalting was 150 ml.
- The desalted sample was applied to a Source Q30 anion ex-35 changer column (100 ml, 5x5 cm) equilibrated in 20 mM triethanolamine, pH 7.3. The column was washed with equilibration buffer until a stable baseline was obtained.

24 Lipase activity was eluted with a 420 ml linear gradient from 0 to 0.35 M sodium chloride in equilibration buffer, flow rate 5 ml/min. Fractions of 10 ml were collected. Sodium acetate (100 µl of a 2M solution) was added to each fraction 5 to adjust pH to 5.5. Fractions 26-32 (70 ml) were pooled. To the pool from the anion exchange step was added ammonium sulfate to 1 M and the sample was applied to a Source Phenyl HIC column (20 ml, 10x2 cm) equilibrated in 20 mM sodium ace-10 tate (pH 5.5), 1 M ammonium sulfate. The column was washed with the equilibration buffer. Lipase was eluted with a 320 ml linear gradient from 1 M to 0 M ammonium sulfate in 20 mM sodium acetate (pH 5.5), flow 1.5 ml/min. Fractions of 7.5 ml were collected. 15 Fractions 33-41 were analyzed by SDS-PAGE using a NOVEX system with precast gels. Both electrophoresis and silver staining of the gels were done according to the manufacturer (Novex, San Diego, USA). (The same system was used for native 20 electrophoresis and isoelectric focusing). It was found that fraction 40 and 41 contained lipase as the only protein. 2.3. Characterization of the purified lipase 25 (i) Determination of molecular weight The apparent molecular weight of the native lipase was 37.7 kDa as measured by the above SDS-PAGE procedure. The purified lipase eluted at a molecular weight of 32.2 kDa from a Supe-30 rose 12 gel filtration column (50 mM sodium phosphate, 0.2 M sodium chloride, pH 6.85, flow 0.65 ml/min) and is therefore a monomer. The molecular weight of the lipase was also determined by 35 matrix-assisted laser desorption ionisation (MALDI) by means of a time-of-flight (TOF) mass spectrometer (Voyager Bio-Spectrometry Workstation, Perspective Biosystems). Samples were prepared by mixing 0.7 µl of desalted lipase solution

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7 μl of a matrix solution containing sinoxy-4-hydroxy cinnamic acid) in 70% ace

and 0.7 µl of a matrix solution containing sinapic acid (3.5-dimethoxy-4-hydroxy cinnamic acid) in 70% acetonitrile (0.1% TFA, 10 mg/ml). 0.7 µl of the sample mixture was placed on top of a stainless steel probe tip and allowed to air-dry prior to introduction into the mass spectrometer. Spectra were obtained from at least 100 laser shots and averaged to obtain a good signal to noise ratio. The molecular mass for the lipase was found to be 30,384 Da and 30,310 Da by two independent analyses.

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Digestion of the lipase with endo-β-N-acetyl-glucosamidase H (10 μl) from Streptomyces (Sigma) was carried out by adding 200 μl lipase and incubating at 37°C for 2 hours. The digestion mixture was desalted using a VSWP filter and analyzed directly by MALDI mass spectrometry. A major component of deglycosylated lipase gave a mass of 29,339 Da and 29,333 Da by two independent analyses. A minor component with a mass of 29,508 Da was also observed. These values corresponds well to the later calculated theoretical value of 28,939 Da based on 20 the complete amino acid sequence of the mature lipase.

#### (ii) Determination of the isoelectric point

The isoelectric point (pI) for the lipase was determined by 25 isoelectric focusing and was found to be 4.1.

A calculation of the pI based on the amino acid sequence as determined in the following and shown as SEQ ID NO: 9 gave an estimated pI of 4.07.

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#### (iii) Determination of temperature stability

Eppendorf tubes with 25  $\mu$ l of purified lipase 3 plus 50  $\mu$ l 100 mM sodium acetate buffer (pH 5.0) were incubated for 1 35 hour in a water bath at respectively 30, 40, 50, and 60°C. A control was treated in the same way, but left at room temperature. After 1 hour the lipase 3 activity was determined by the p-nitrophenyl acetate assay as described above.

The purified lipase had a good thermostability. It was found that the lipase maintained 60% of its activity after 1 hour at 60°C. 80% and 85% activity was maintained after 1 hour at 50°C and 40°C respectively.

#### (iv) Determination of pH stability

Purified lipase 3 (200 µl) was added to 5 ml of 50 mM buffer solutions: (sodium phosphate, pH 8.0, 7.0 and 6.0 and sodium acetate pH 5.0, 4.0 and 3.5). The control was diluted in 5 ml of 4 mM sodium acetate pH 5.5. After four days at room temperature the residual activity was measured by the Modified Food Chemical Codex assay for lipase activity as described above. The lipase was very stable in the pH range from 4.0 to 7.0 where it maintained about 100% activity relative to the control (Table 2.1). At pH 3.5 the lipase maintained 92% activity, and at pH 8.0 95% residual activity was maintained as compared to the control.

Table 2.1. pH stability of lipase 3

рН	Activity (LUT/ml)	Activity (%)
Control (pH 5.5)	89.2	100
3.5	82.5	92
4.0	91.7	103
5.0	86.5	97
6.0	92.4	104
7.0	90.6	102
8.0	84.4	95

20

EXAMPLE 3

### Amino acid sequencing of lipase 3

- 5 Purified lipase enzyme was freeze-dried and 100  $\mu g$  of the freeze-dried material was dissolved in 50  $\mu l$  of a mixture of 8 M urea and 0.4 M ammonium hydrogencarbonate, pH 8.4. The dissolved protein was denatured and reduced for 15 minutes at 50°C following overlay with nitrogen and addition of 5  $\mu l$  45
- 10 mM dithiothreitol. After cooling to room temperature, 5  $\mu l$  of 100 mM iodoacetamide was added for the cysteine residues to be derivatized for 15 minutes at room temperature in the dark under nitrogen.
- 15 135  $\mu$ l of water and 5  $\mu$ g of endoproteinase Lys-C in 5  $\mu$ l of water was added to the above reaction mixture and the digestion was carried out at 37°C under nitrogen for 24 hours. The resulting peptides were separated by reverse phase HPLC on a VYDAC C18 column (0.46  $_{\times}$  15 cm; 10  $\mu$ m; The Separation
- 20 Group, California, USA) using solvent A: 0.1% TFA in water and solvent B: 0.1% TFA in acetonitrile. Selected peptides were rechromatographed on a Develosil C18 column (0.46  $\times$  10 cm, Novo Nordisk, Bagsværd, Denmark) using the same solvent system, prior to N-terminal sequencing. Sequencing was done
- 25 using an Applied Biosystems 476A sequencer using pulsedliquid fast cycles according to the manufacturer's instructions (Applied Biosystems, California, USA).

For direct N-terminal sequencing, the purified protein was 30 passed through a Brownlee C2 Aquapore column (0.46  $\times$  3 cm, 7  $\mu$ m, Applied Biosystems, California, USA) using the same solvent system as above. N-terminal sequencing was then performed as described above. As the protein was not derivatized prior to sequencing, cysteine residues could not be 35 determined.

The following peptide sequences were found:

N-terminal:

Ser-Val-Ser-Thr-Ser-Thr-Leu-Asp-Glu-Leu-Gln-Leu-Phe-Ala-Gln-Trp-Ser-Ala-Ala-Ala-Tyr-X-Ser-Asn-Asn

(SEQ ID NO:1)

5

Internal peptide 2: Ala-Trp-Glu-Ser-Ala-Ala-Asp-Glu-Leu
Thr-Ser-Lys-Ile-Lys
(SEQ ID NO:3)

No further peptides could be purified from the HPLC fractionation presumably because they were very hydrophobic and therefore tightly bound to the reverse phase column.

A search in SWISS-PROT database release 31 for amino acid sequences with homology to the above peptides was performed and only three sequences were found.

20

All of the above peptides showed a low homology to the above known sequences. Especially internal peptide 2 has very low homology to the three lipases, LIP-RHIDL, LIP-RHIMI and MDLA-PENCA from *Rhizopus delamar* (Haas and Berka, Gene, 1991,

25 109:107-113), Rhizomucor miehei (Boel et al., Lipids, 1988, 23:701-706) and Penicillium camenbertii (Yamaguchi et al., Gene, 1991, 103:61-67; Isobe and Nokihara, Febs. Lett., 1993, 320:101-106) respectively. Although the homology was not very high it was possible to position the lipase 3 peptides on 30 these sequences as it is shown in the below Table 3.1.



# Table 3.1. Alignment of lipase 3 peptides with known lipase sequences

35	LIP_RHIDL	MVSFISISQGVSLCLUSSMMLGSSAVPVSGKSGSSNTAVSASDNAALPP	50
	LIP_RHIMI	MVLKQRANYLGFLIVFTAFLVEAVPIKRQSNSTVDSLPP	4 C
	MDLA_PENCA	MRLSSAVASLGYALPG	21

SVSTSTLDELOLFAOWSAAAYXSNN N-Terminal LIP RHIDL LISSRCAPPSNKGSKSDLOAEPYNMOKNTEWYESHGGNLTSIGKRDDNLV 100 LIP RHIMI LIPSRTSAPSSSPSTTDPEAPAM-----SRNGPLPS----DVETK 76 5 MDLA PENCA KLQSR----~-DVSTSELDQFEFWVQYAAASY----------47 LIP RHIDL GGMTLDLPSDAPPISLSSTNSASDGGKVVAATTAOIOEFTKYAGIAATA 150 LIP RHIMI 121 YGMALNATSYPDSV----VQAMSIDGGIRAATSQEINELTYYTTLSANS 10 MDLA PENCA -----YEADYTAOVGDKL 60 LIP RHIDL YCRSVVPGNKWDCVQCQKWVPDGKIITTFT-SLLSDTNGYVLRSDKQKTI 199 LIP RHIMI YCRTVIPGATWDCIHCDA-TEDLKIIKTWS-TLIYDTNAMVARGDSEKTI 169 15 MDLA PENCA SCSKG----NCPEVEA--TGATVSYDFSDSTITDTAGYIAVDHTNSAV ... . . \*\*.. . ...... . . . Peptide 1 VHTGFWK Peptide 2 AWESAADELTSK 20 LIP RHIDL YLVFRGTNSFRSAITDIVFNFSDYKPVKGAKVHAGFLSSYEQVVNDYFPV 249 LIP RHIMI YIVFRGSSSIRNWIADLTFVPVSYPPVSGTKVHKGFLDSYGEVQNELVAT 219 MDLA PENCA VLAFRGSYSVRNWVADATFVHTNPGLCDGCLAELGFWSSWKLVRDDIIKE 152 25 Peptide 2 ΙK LIP RHIDL VQEQLTAHPTYKVIVTGHSLGGAQALLAGMDLYQREPRLSPKNLSIFTVG 299 LIP RHIMI VLDQFKQYPSYKVAVTGHSLGGATALLCALDLYQREEGLSSSNLFLYTQG 269 30 MDLA PENCA LKEVVAQNPNYELVVVGHSLGAAVATLAATDL--RGKGYPSAKLYAYA--198 LIP RHIDL GPRVGNPTFAYYVESTGIPFORTVHKRDIVPHVPPQSFGFLHPGVESWIK 349 LIP RHIMI QPRVGDPAFANYVVSTGIPYRRTVNERDIVPHLPPAAFGFLHAGEEYWIT 319 35 MDLA PENCA SPRVGNAALAKYITAQGNNF-RFTHTNDPVPKLPLLSMGYVHVSPEYWIT 247 \*\*\*\*...\* \*. . \* . . \* \*\*..\* ..\*..\* . \* \*\*. LIP RHIDL SGTSN-V----QICTSEIETKDCSNSIVPFTSILD-HLSYF-DINEGSC 391 LIP RHIMI DNSPETV----QVCTSDLETSDCSNSIVPFTSVLD-HLSYF-GINTGLC 362 40 MDLA PENCA 297 SPNNATVSTSDIKVIDGDVSFDGNTGTGLPLLTDFEAHIWYFVQVDAGKG LIP\_RHIDL ----L 392 LIP RHIMI ----T 363 45 MDLA PENCA PGLPFKRV 305

#### EXAMPLE 4

# Isolation and purification of Aspergillus tubigensis genomic DNA

5

The Aspergillus tubigensis mutant strain was grown in PDB (Difco) for 72 hours and the mycelium was harvested. 0.5-1 g of mycelium was frozen in liquid nitrogen and ground in a mortar. Following evaporation of the nitrogen, the ground 10 mycelium was mixed with 15 ml of an extraction buffer (100 mM

- Tris·HCl, pH 8.0, 50 mM EDTA, 500 mM NaCl, 10 mM ß-mercaptoe-thanol) and 1 ml 20 % sodium dodecylsulfate. The mixture was vigorously mixed and incubated at 65°C for 10 min. 5 ml 3M potassium acetate, (pH 5.1 adjusted with glacial acetic acid)
- 15 was added and the mixture further incubated on ice for 20 min. The cellular debris was removed by centrifugation for 20 min. at 20,000  $\times$  g and 10 ml isopropanol was added to the supernatant to precipitate (30 min at -20°C) the extracted DNA. After further centrifugation for 15 min at 20,000  $\times$  g,
- 20 the DNA pellet was dissolved in 1 ml TE (10 mM Tris·HCl pH 8.0, 1 mM EDTA) and precipitated again by addition of 0.1 ml 3 M NaAc, pH 4.8 and 2.5 ml ethanol. After centrifugation for 15 min at 20,000  $_{\rm X}$  g the DNA pellet was washed with 1 ml 70 % ethanol and dried under vacuum. Finally, the DNA was dis-
- 25 solved in 200  $\mu$ l TE and stored at -20°C.

#### EXAMPLE 5

# The generation of a fragment of the putative gene coding for lipase 3 using PCR

To obtain a fragment of the putative gene (in the following referred to as the *lipA* gene) as a tag to isolate the complete gene, a PCR amplification procedure based on the infomation in the isolated peptide sequences was carried out.

3 JAN

Degenerated primers for RCR amplification of a fragment of

the lipase gene were designed based on the amino acid sequences of the isolated peptides. The following three PCR primers were synthesised:

5 CO35: TTC CAR AAN CCN GTR TGN AC (SEQ ID NO:4)

20 mer 256 mixture, based on peptide 1 sequence VHTGFWK (Reversed).

10 C036: CAR YTN TTY GCN CAR TGG (SEQ ID NO:5)

18 mer 256 mixture, based on the N-terminal sequence QLFAQW.

C037: GCV GCH SWY TCC CAV GC (SEQ ID NO:6)

17 mer 216 mixture, based on internal peptide 2 sequence AWESAA (reversed).

The oligonucleotides were synthesised on a Applied Biosystems 20 model 392 DNA/RNA Synthesizer. To reduce the degree of degeneracy the rare Ala codon GCA and the Ser codon TCA have been excluded in design of primer CO37.

With these primers the desired fragments were amplified by 25 PCR. Using these primers it was expected that a fragment of about 300 bp should be amplified provided there are no introns in the fragment.

The following PCR reactions were set up in 0.5 ml PCR tubes 30 to amplify a putative lipA fragment:

1. 0.5  $\mu$ g total genomic DNA,

100 pmol primer C036,

100 pmol primer C037,

35 10 μl PCR Buffer II (Perkin Elmer),

6 µl 25 mM MgCl<sub>2</sub>,

2 µl dNTP mix (10 mM dATP, 10 mM dCTP, 10 mM dGTP, 10 mM

dTT

2 units Amplitaq polymerase (Perkin Elmer), and

water to a total volume of 100  $\mu$ l.

2. 0.5  $\mu$ g total genomic DNA,

100 pmol primer C035,

5 100 pmol primer C036,

10 µl PCR Buffer II (Perkin Elmer),

6  $\mu$ l 25 mM MgCl<sub>2</sub>,

2 µl dNTP mix (10 mM dATP, 10 mM dCTP, 10 mM dGTP, 10 mM

dTT

2 units Amplitag polymerase (Perkin Elmer), and

10 water to a total volume of 100 μl.

The reactions were performed using the following program:

94°C 2 min

94°C 1 min )

15 40°C 1 min )

72°C 1 min ) These three steps were repeated for 30

72°C 5 min cycles

5°C SOAK

20 The PCR amplifications were performed in a MJ Research Inc. PTC-100 Thermocycler.

In reaction 1, three distinct bands of about 300, 360 and 400 bp, respectively could be detected. These bands were isolated

- 25 and cloned using the pT7-Blue-T-vector kit (Novagene). The sizes of these fragment is in agreement with the expected size provided that the fragment contains 0, 1 or 2 introns, respectively.
- 30 The three fragments were sequenced using a "Thermo Sekvenase fluorescent labelled primer cycle sequencing Kit" (Amersham) and analyzed on a ALF sequencer (Pharmacia) according to the instructions of the manufacturer. The fragment of about 360 bp contained a sequence that was identified as a lipase and,
- 35 as it contained the part of the N-terminal distal to the sequence used for primer design, it was concluded that the desired *lip*A gene fragment was obtained.

The sequence of the about 360 bp PCR fragment (SEQ ID NO:7) is shown in the following Table 5.1. The peptide sequence used for primer design is underlined. The remaining part of 5 the N-terminal sequence is doubly underlined.

3ut	Table	: :	5.1	1.\	PC:	R-ç	en	era	te	d p	ut	ati	ve	1i	pΑ	se	que	enc	<u>e</u>		
*/				\																	
					10			20			3	0			40			50			60
10					1			ŀ				1			1			1			1
	t	tacccggggntccgatt <u>CAGTTGTTCGCGCAATGG</u> TCTGCCGCAGCTTATTGCTCGAAT														ATA					
	Q L F A Q W S A A A Y C S													<u>S</u>	<u>N</u>						
1 -					7.0						^	^		•	00			110			120
15					70	1		80			9	0		1	1			110			120
	7	מית	ጥር	CAC	ነ ጥርር	<b>ά</b> αα:	GAV	ו ייירר	ממ	ጥጥር	מרמ	ו דכר	ACG	GCC	ו רום ב	3CC,	гст	CCA	тса	GTC	GAGG
	ATATCGACTCGAAAGAVTCCAACTTGACATGCACGGCCAACGCCTGTCCATCAGTCGAGG																				
		N	I	D	S	K	`\ <b>x</b>	S	N	L	т	С	Т	А	N	Α	С	Р	s	v	E
20	-																				
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	A	GG	CCZ	AGT	ACC	ACG	ATG	дтс	CTG	GAG	TTC	GAC	CTG	TAT	GTC	ACT	CAG	ATC	GCA	GAC.	ATAG
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25	E	;	A	s	T	T	M	$\mathbf{r}_{j}^{'}$	L	E	F	D	L	Y	V	T	Q	I	Α	D	I
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				1	90			200			210 220					230					240
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30	A	.GC	AC	AGC	TAA	TTG	AAC	<b>AG</b> G	ACG	AAC	GAC	TTT	TGG	AGG	CAC.	AGC	-66	111	CCI	GGC	CGCG
30	E		Н	s	_	L	N	R	T	N	D	F	W	R	н	s	R	F	P	G	R
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				2	50			260		/	270 280 290									300	
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35	G	AC	AA	CAC	CAA	.CAA	.GCG	GCT	CGT	GGT	CGC	СТТ	CCG	GGG	AAG	CAG	CAC	GAT	TGA	GAA	CTGG
	G	;	Q	Н	Q	Q	A	A	R	G	R	L	P	G	K	Q	Н	D	-	E	L
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40	O I I I I I I I I I I I I I I I I I I I																				
	A	тт	GC:	ΓAΑ	TCY	TGA	CTT	CAT	CCT	GGR	AGA	TAA,	.CG								
	_		_		.,		7	,,		v	_	•									
	D	1	C	_	Х	_	Т	Н	Р	Х	K	7.									

The finding of this sequence permitted full identification of the PCR fragment as part of the lipA gene. The stop codon found in the reading frame can be caused either by a PCR or a reading error or there can be an intron encoded in the fragment as a consensus intron start and ending signal (shown in bold). If the putative intron is removed a shift in reading frame will occur. However, an alignment of the deduced amino acid sequence and the fungal lipases shown in Table 3.1 suggested that the fragment was part of the desired gene.

10

EXAMPLE 6

### Cloning and characterisation of the lipA gene

#### 15 (i) Construction of an Aspergillus tubigensis genomic library

Aspergillus tubigensis genomic DNA was digested partially with Tsp5091 (New England Biolabs Inc.). 10  $\mu$ g DNA was digested in 100  $\mu$ l reaction mixture containing 2 units Tsp5091.

- 20 After 5, 10, 15 and 20 minutes 25  $\mu$ l was removed from the reaction mixture and the digestion was stopped by addition of 1  $\mu$ l 0.5 M EDTA, pH 8.0. After all four reactions had been stopped, the samples were run on a 1% agarose gel in TAE buffer (10 x TAE stock containing per litre: 48.4 g Trizma
- 25 base, 11.5 ml glacial acetic acid, 20 ml 0.5 M EDTA pH 8.0).

  HindIII-digested phage Lambda DNA was used as molecular

  weight marker (DNA molecular weight marker II, Boehringer,

  Mannheim). Fragments of a size between about 5 and 10 kb were

  cut out of the gel and the DNA fragments were purified using
- 30 Gene Clean II Kit (Bio-101 Inc.). The purified fragments were pooled and 100 ng of the pooled fragments were ligated into 1  $\mu$ g EcoRI-digested and dephosphorylated ZAP II vector (Stratagene) in a total volume of 5  $\mu$ l. 2  $\mu$ l of this volume was packed with Gigapack II packing extract (Stratagene) which 35 gave a primary library of 650,000 pfu.
  - $E.\ coli$  strain XL1-Blue-MRF (Stratagene) was infected with 5x 50,000 pfu of the primary library. The infected bacteria were

35 mixed with top agarose (as NZY plates but with 6 g agarose per litre instead of the agar) and plated on 5 NZY plates (13 cm). After incubation at 37°C for 7 hours, 10 ml SM buffer (per litre: 5.8 g NaCl, 2.0 g MgCl<sub>2</sub> 7H<sub>2</sub>O, 50 ml 1 M Tris HCl 5 pH 7.5, 5.0 ml of 2% (w/v) gelatine) and incubated overnight at room temperature with gently shaking. The buffer containing washed-out phages was collected and pooled. 5% chloroform was added and after vigorous mixing the mixture was incubated 1 hour at room temperature. After centrifugation for 2 min-10 utes at  $10,000 \times g$  the upper phase containing the amplified library was collected and dimethylsulphoxide was added to 7%. Aliquots of the library was taken out in small tubes and frozen at -80°C. The frozen library contained 2.7  $\times$  10 $^9$ pfu/ml with about 6% without inserts. 15 (ii) Screening of the Aspergillus tubiqensis library  $2 \times 50.000$  pfu were plated on large (22 x 22 cm) NZY plates containing a medium containing per litre: 5 g NaCl, 2 g 20 MgSO<sub>4</sub>·7H<sub>2</sub>O, 5 g yeast extract, 10 g casein hydrolysate, 15 g agar, pH adjusted to 7.5 with NaOH. The medium was autoclaved and cooled to about 60°C and poured into the plates. Per plate was used 240 ml of medium. 25 The inoculated NZY plates were incubated overnight at 37°C and plaque lifts of the plates were made. Two lifts were made for each plate on Hybond N (Amersham) filters. The DNA was fixed using UV radiation for 3 min. and the filters were hybridized as described in the following using, as the probe, 30 the above PCR fragment of about 360 bp that was labelled with <sup>32</sup>P-dCTP using Ready-to-Go labelling kit (Pharmacia). The filters were prehybridised for one hour at 65°C in 25 ml

prehybridisation buffer containing 6.25 ml 20 x SSC (0.3 M 35 Na<sub>3</sub>citrate, 3 M NaCl), 1,25 ml 100 x Denhard solution, 1.25 ml 10% SDS and 16.25 ml water. 150 µl 10 mg/ml denatured Salmon sperm DNA was added to the prehybridization buffer

36 immediately before use. Following prehybridization, the prehybridisation buffer was discarded and the filters hybridised overnight at 65°C in 25 ml prehybridisation buffer with the radiolabelled PCR fragment. 5 Next day the filters were washed according to the following procedure:  $2 \times 15$  min. with  $2 \times SSC + 0.1 \% SDS, 15$  min. with 1 x SSC + 0.1 % SDS and 10 min. with 0.1 x SSC + 0.1% SDS. 10 All washes were done at 65°C. The sheets were autoradiographed for 16 hours and positive clones were isolated. A clone was reckoned as positive only if there was a hybridisation signal on both plaque lifts of the plate in question. 15 Seven putative clones were isolated and four were purified by plating on small petri dishes and performing plaque lifts essentially as described above. The purified clones were converted to plasmids using an 20 ExAssist Kit (Stratagene). Two sequencing primers were designed based on the about 360 bp PCR fragment. The sequencing primers were used to sequence the clones and a positive clone with the lipA gene encoding 25 lipase 3 was found. The isolated positive clone was designated pLIP4. (iii) Characterisation of the pLIP4 clone 30 A restriction map of the clone was made. The above 360 bp PCR fragment contained a SacII site and as this site could be found in the genomic clone as well this site facilitated the construction of the map. The restriction map showing the structure of pLIP4 is shown in Fig. 1. The restriction map 35 shows that the complete gene is present in the clone. Additionally, since promoter and terminator sequences are present, it was assumed that all the important regions is present in the clone.

A sample of Escherichia coli strain DH5α containing pLIP4 was deposited in accordance with the Budapest Treaty with The 5 National Collections of Industrial and Marine Bacteria Limited (NCIMB) at 23 St. Machar Drive, Aberdeen, Scotland, United Kingdom, AB2 1RY on 24 February 1997 under the accession number NCIMB 40863.

- 10 The gene was sequenced using cycle sequencing and conventional sequencing technology. The complete sequence (SEQ ID NO:8) is shown below in Table 6.1. The sequence has been determined for both strands for the complete coding region and about 100 b $\dot{p}$  upstream and downstream of the coding
- 15 region. The sequences downstream to the coding region have only been determined on one strand and contains a few uncertainties. In the sequence as shown below, the intron sequences are indicated as lowercase letters and the N-terminal and the two internal peptides (peptide 1 and peptide 2)
- 20 are underlined:

### Table 6.1. The DNA sequence for the lipA gene and flanking sequences

- 25 1 CCNDTTAATCCCCCACCGGGGTTCCCGCTCCCGGATGGAGATGGGGCCAAAACTGGCAAC
  - CCCCAGTTGCGCAACGGAACAACCGCCGACCCGGAACAAAGGATGCGGATGAGGAGATAC
  - 121 GGTGCCTGATTGCATGGCTGGCTTCATCTGCTATCGTGACAGTGCTCTTTGGGTGAATAT
  - 181 TGTTGTCTGACTTACCCCGCTTCTTGCTTTTTCCCCCCTGAGGCCCTGATGGGGAATCGC
  - 241 GGTGGGTAATATGATATGGGTATAAAAGGGAGATCGGAGGTGCAGTTGGATTGAGGCAGT
- 30 301 GTGTGTGTGCATTGCAGAAGCCCGTTGGTCGCAAGGTTTTGGTCGCCTCGATTGTTTG
  - 361 TATACCGCAAGATGTTCTCTGGACGGTTTGGAGTGCTTTTGACAGCGCTTGCTGCCTGG
    - MFSGRFGVLLTALAAL
  - 421 GTGCTGCCGCGCCGCACCGCTTGCTGTGCGGAgtaggtgtgcccgatgtgagatggttg GAAAPAPLAVR
- 35 481 gatagcactgatgaagggtgaatagGTGTCTCGACTTCCACGTTGGATGAGTTGCAATTG S V S T S T L D E L Q L
  - 541 TTCGCGCAATGGTCTGCCGCAGCTTATTGCTCGAATAATATCGACTCGAAAGACTCCAAC FAQWSAAAYC\$NNIDSKDSN
  - 601 TTGACATGCACGGCCAACGCCTGTCCATCAGTCGAGGAGGCCAGTACCACGATGCTGCTG
- 40 L T C T A N A C P S V E E A S T T M L L
  - 661 GAGTTCGACCTgtatgtcactcagatcgcagacatagagcacagctaatttgaacagGAC

E F D L GAACGACTTTGGAGGCACAGCCGGTTTCCTGGCCGCGGACAACACCAACAAGCGGCTCGT N D F G G T A G F L A A D N T N K R L V GGTCGCCTTCCGGGGAAGCAGCACGATTGAGAACTGGATTGCTAATCTTGACTTCATCCT V A F R G S S T I E N W I A N L GGAAGATAACGACGACCTCTGCACCGGCTGCAAGGTCCATACTGGTTTCTGGAAGGCATG N D D L C T G C K V H T G F W K A W GGAGTCCGCTGCCGACGAACTGACGAGCAAGATCAAGTCTGCGATGAGCACGTATTCGGG 901 E S A A D \E L T S K I K S A M S T Y S G 10 961 CTATACCCTATACTTCACCGGGCACAGTTTGGGCGGCGCATTGGCTACGCTGGGAGCGAC Y T L Y F T G H S L G G A L A T L G A T 1021 AGTTCTGCGAAATGACGGATATAGCGTTGAGCTGgtgagtccttcacaaaggtgatggag V L R N D G \ Y S V E L 1081 cgacaatcgggaacagacadtcaatagTACACCTATGGATGTCCTCGAATCGGAAACTAT YTYGCPRIGNY 15 1141 GCGCTGGCTGAGCATATCACCAGTCAGGGATCTGGGGCCAACTTCCGTGTTACACACTTG A L A E H I T\S Q G S G A N F R V T H L N D I V P R V P P M D F G F S Q P S P E 20 1261 TACTGGATCACCAGTGGCAATGGAGCCAGTGTCACGGCGTCGGATATCGAAGTCATCGAG I T S G N G \A S V T A S D I E V I E 1321 GGAATCAATTCAACGGCGGGAAATGCAGGCGAAGCAACGGTGAGCGTTGTGGCTCACTTG G I N S T A G N Å G E A T V S V V A H L 25 1381 TGGTACTTTTTTGCGATTTCCGAGTGCCTGCTATAACTAGACCGACTGTCAGATTAGTGG WYFFAISECLLL 1441 ACGGGAGAGTGTACATAAGTAATTAGTATAATCAGAGCAACCCAGTGGTGGTGATGG 1501 TGGTGAAAGAACACATTGAGTTCCCATTACGKAGCAGWTAAAGCACKTKKGGAGGC 1561 GCTGGTTCCTCCACTTGGCAGTTGGCGGCCATCAATCATCTTTCCTCTCCTTACTTTCGT 30 1621 CCACCACAACTCCCATCCTGCCAGCTGTCGCATCCCCGGGTTGCAACAACTATCGCCTCC 1681 GGGGCCTCCGTGGTTCTCCTATATTATTCCATCCGACGCCGACGTTTCACCCTCAACCT 1801 TCACGACCCGACCGTCTGYGATYGTCCAACCG

### 35 (iv) Analysis of the sequence of the complete gene

The peptide sequences obtained could all be found in the deduced amino acid sequence (see Table 5.1) which confirms again that the sequence found is the sequence of the lipase 3 40 gene. The gene was designated *lipA*.

The amino acid sequence was aligned with the three fungal lipases used to align the peptide sequences. The alignment is shown in Table 6.2.

# Table 6.2 Alignment of the lipase 3 sequence with known fungal lipases

5	LIPASE3	MFSGTALAA	15
	MDLA_PENCA	MRLSSAVAS	14
	LIP_RHIDL	MVSFISISQGVSLCLLVSSMMLGSSAVPVSGKSGSSNTAVSASDNAALPP	50
	LIP_RHIMI	MVLKQRANYLGFLIVFFTAFLVEAVPIKRQSNSTVDSLPP	40
		*	
10		i i	
	LIPASE3	L	16
	MDLA_PENCA	L	15
	LIP_RHIDL	LISSRCAPPSNKGSKSDLQAEPYNMQKNTEWYESHGGNLTSIGKRDDNLV	100
	LIP_RHIMI	LIPSRTSAPSSSPSTTDPEAPAMSRNGPLPSDVETK	76
15		*	
		AND CHOMOME DELOT ENOUGANA	47
	LIPASE3	GAAAPAPLAVRSVSTSTLDELQLFAQWSAAA	4 6
	MDLA_PENCA	GYALPGKLQSRDVSTSELDQFEFWVQYAAAS	150
20	LIP_RHIDL	GGMTLDLPSDAPPISL\$SSTNSASDGGKVVAATTAQIQEFTKYAGIAATA	121
20	LIP_RHIMI	YGMALNATSYPDSVVQAMSIDGGIRAATSQEINELTYYTTLSANS	121
		• • • • • • • • • • • • • • • • • • • •	
	LIPASE3	YCSNNIDSK-DSNLTCTANACPSVEEASTTMLLEFDLTNDFGGTAGFLAA	96
	MDLA PENCA	YYEADYTAQVGDKLSCSKGNCPEVEATGATVSYDFS-DSTITDTAGYIAV	95
25	LIP RHIDL	YCRSVVPGNKWDCVQCQKWVPDGKIITTFTSLLSDTNGYVLR	192
23	LIP RHIMI	YCRTVIPGATWDCIHCDA-TEDLKIIKTWSTLIYDTNAMVAR	162
	227	* * *	
	LIPASE3	DNTNKRLVVAFRGSSTIENWI <b>A</b> NLDFILEDNDDLCTGCKVHTGFWKAWES	146
30	MDLA PENCA	DHTNSAVVLAFRGSYSVRNWVADATFV-HTNPGLCDGCLAELGFWSSWKL	144
	LIP_RHIDL	SDKQKTIYLVFRGTNSFRSAITDIVFNFSDYKPV-KGAKVHAGFLSSYEQ	241
	LIP_RHIMI	GDSEKTIYIVFRGSSSIRNWIADLTFVPVSYPPV-SGTKVHKGFLDSYGE	211
		***	
35	LIPASE3	AADELTSKIKSAMSTYSGYTLYFTGHSLGGALATLGATVLRNDGY-SV	193
	_	VRDDIIKELKEVVAQNPNYELVVVGHSLGAAVATLAATDLRGKGYPSA	192
	LIP_RHIDL	VVNDYFPVVQEQLTAHPTYKVIVTGHSLGGAQALLAGMDLYQREPRLSPK	291
	LIP_RHIMI	VQNELVATVLDQFKQYPSYKVAVTGHSLGGATALLCALDLYQREEGLSSS	261
40	<b>_</b>		2 4 1
	LIPASE3	ELYTYGCPRIGNYALAEHITSQGSGANFRVTHLNDIVPRVPPMDFGFS	241 238
		KLYAYASPRVGNAALAKYITAQGNNFRFTHTNDPVPKLPLLSMGYV	340
	LIP_RHIDL	NLSIFTVGGPRVGNPTFAYYVESTGIPFQ-RTVHKRDIVPHVPPQSFGFL	310
4 5	LIP_RHIMI	NLFLYTQGQPRVGDPAFANYVVSTGIPYR-RTVNERDIVPHLPPAAFGFL	210
45		.* **.** * * ****.	

*						
Com 10	LIPASE3	QPSREYWITSGNGASVTAS	DIEVIEGINSTAGN	AGEATVSVV	AHLWY	288
' <b>'</b>	MDLA_PENCA	HVSPEYWITSPNNATVSTS	DIKVIDGDVSFDGN	TGTGLPLLTD	FEAHIWY	288
A Comment	LIP_RHIDL	HPGVESWIKSGTSN-VQIC	TSEIETKD	CSNSIVPFTS	ILDHLSY	383
( 5	LIP_RHIMI	HAGEEYWÎTDNSPETVQVC	TSDLETSD	CSNSIVPFTSV	<b>VLDHLSY</b>	354
		* ** * .		• • •	.*. *	
	LIPASE3	FFAISECLL	297			
	MDLA_PENCA	fvqvdagkgpgl <b>a</b> fkrv	305			
10	LIP_RHIDL	F-DINEGSC	392			
	LIP_RHIMI	F-GINTGLC	363			

The above alignment shows that lipase 3 is homologous to the 15 known lipase sequences but that the homology is not very high. Deletions or insertions in the lipase 3 sequence was not observed when comparing the sequence with these three lipases. This strengthens the probability that the putative introns have been identified correctly.

20

A search in SWISS-PROT release 31 database was performed and it did not lead to further sequences with higher homology than that to the above known lipases (Table 6.3).

25 The sequence with highest homology is a mono- diacyl lipase from *Penicillium camembertii* where the identity is found to 42 %. However the C- terminal of lipase 3 resembles the 2 lipases from Zygomycetes (*Rhizopus* and *Rhizomucor*) and not the *P. camembertii* enzyme.

30

# Table 6.3. Alignment of coding sequences of the lipA gene and gene coding for mono-diacyl lipase from Penicillium camemberti

- NKRLVVAFRGSSTIENWIANLDFILEDNDDLCTGCKVHTGFWKAWESAAD -149 LIPASE3 11 1 1 1 11 11 111 T IIIÀE 5 MDLA\_PENCA- NSAVVLAFRG\$YSVRNWVADATFV-HTNPGLCDGCLAELGFWSSWKLVRD -147 - ELTSKIKSAMSTYSGYTLYFTGHSLGGALATLGATVLRNDGY-SVELYTY -198 LIPASE3 11111 1 11 11 11 11 11 1 MDLA PENCA- DIIKELKEVVAQNPNYELVVVGHSLGAAVATLAATDLRGKGYPSAKLYAY -197 10 - GCPRIGNYALAEHITSQGSGANFRVTHLNDIVPRVPPMDFGFSQPSPEYW -248 LIPASE3 11 11 111 111 MDLA\_PENCA- ASPRVGNAALAKYİTAQGN--NFRFTHTNDPVPKLPLLSMGYVHVSPEYW -245 - ITSGNGASVTASDIEVIEGINSTAGNAGEATVSVV---AHLWYFFAISEC -295 15 LIPASE3 11 111 MDLA\_PENCA- ITSPNNATVSTSDIKVIDGDVSFDGNTGTGLPLLTDFEAHIWYFVQVDAG -295 - L----L -297 LIPASE3 20 MDLA PENCA- KGPGLPFKRV -305 Identity: 126 amino acids (42.42%)

25 The N-terminal of the mature lipase has been determined by N-terminal sequencing to be the serine residue No. 28 of the lipase 3 precursor (SEQ ID NO:9) as shown in Table 6.4 below. Hence the amino acids No. 1 to No. 27 is the signal sequence.

Amino acid sequence of the precursor of lipase 3 30 Table 6.4: 20 25 30 10 1 M F S G R F G V L L T A L A A L G A A A P A P L A V R S V S 31 TSTLD L LQLFAQWSAAAYCSNNIDSKDSNL 61 T C T A N A C P S V E E A S T T M L L E F D L T N D F G G T 35 91 A G F L A A\D N T N K R L V V A F R G S S T I E N W I A N L 121 D F I L E D N D D L C T G C K V H T G F W K A W E S A A D E 151 L T S K I K S\A M S T Y S G Y T L Y F T G H S L G G A L A T 181 L G A T V L R N D G Y S V E L Y T Y G C P R I G N Y A L A E 211 H I T S Q G S G\A N F R V T H L N D I V P R V P P M D F G F 40 241 S Q P S P E Y W T T S G N G A S V T A S D I E V I E G I N S 271 TAGNAGEAT VSVVAHLWYFFAISECLL

Number of residues : 297 \

42 Residues 167-176 are recognised as a common motif for the serine lipases (PROSITE). The crystal structure for the Rhizomucor miehei serine lipase has been examined and the 5 residues in the active site identified (Brady et al., Nature, 1990, 343:767-770; Derewanda et al., J. Mol. Biol., 1992, 227:818-839). The active site residues of R. miehei lipase have all been conserved in all the lipases and correspond to the following residues in lipase 3: serine 173, aspartic acid 10 228 and histidine 285. Lipase 3 contains 7 cysteine residues. Four of these are conserved in the P. camembertii lipase where they form disulphide bonds (Isobe and Nokuhara, Gene, 1991, 103:61-67). 15 This corresponds to disulphide bonds between residue 62- 67 and 131-134. In addition, two cysteine residues are homologous to two C residues which forms an additional disulphide bond in Rhizopus and Rhizomucor lipases corresponding to residues 49-295. 20 Two putative N-glycosylation sites were found in lipase 3 in . position 59 and 269. Neither of these are conserved in the other fungal lipases. 25 EXAMPLE 7 Transformation of Aspergillus tubigensis and overexpression of lipase 3 in A. tubigensis 30 The protocol for transformation was based on the teachings of Buxton et al. (Gene, 1985, 37:207-214), Daboussi et al (Curr. Genet., 1989, 15:453-456) and Punt and van den Hondel, (Meth. Enzym., 1992, 216:447-457). 35 A multicopy lipA strain was produced by transforming the pLIP4 plasmid into Aspergillus tubigensis strain 6M 179 using cotransformation with a hygromycin resistant marker plasmid.

45 C. BAKING EXPERIMENTS EXAMPLE 8 5 Baking experiments using lipase 3 8.1. Baking procedures and analytical methods (i) Baking procedure for Danish toast bread 10 Flour (Danish reform flour) 2000 g, dry yeast 30 g, salt 30 g and water corresponding to 400 Brabender units + 3%, was kneaded in a Hobart Mixer with hook for 2 min. at low speed and 10 min. at high speed. Dough temperature after kneading 15 was 25°C. Resting time was 10 min. at 30°C. The dough was scaled 750 g per dough and rested again for 5 min at 33°C and 85% RH. After moulding on a Glimik moulder, the dough were proofed in tins for 50 min at 33°C and baked in a Wachtel oven for 40 min at 220°C with steam injection for 16 sec. 20 After cooling, the bread was scaled and the volume of the bread was measured by the rape seed displacement method. The specific volume is calculated by dividing the bread volume (ml) by the weight (g) of the bread. 25 The crumb was evaluated subjectively using a scale from 1 to 5 where 1 = coarsely inhomogeneous and <math>5 = nicely homogeneous. Three breads baked in tins with lid were stored at 20°C and 30 used for firmness measurements and pore measurements by means of an Image Analyzer. (ii) Baking procedure for Danish rolls 35 Flour (Danish reform) 1500 q, compressed yeast 90 g, sugar 24 g, salt 24 g and water corresponding to 400 Brabender units -2% were kneaded in a Hobart mixer with hook for 2 min. at low speed and 9 min at high speed. After kneading, the dough temperature was 26°C. The dough was scaled 1350 g. After resting for 10 min. at 30°C, the dough was moulded on a Fortuna moulder after which the dough was proofed for 45 min.

5 at 34°C and baked in a Bago oven for 18 min. at 220°C with steam injection for 12 sec. After cooling, the rolls were scaled and the volume of the rolls was measured by the rape seed displacement method. Specific volume is calculated as described above.

10

### (iii) Determination of pore homogeneity

The pore homogeneity of the bread was measured by means of an image analyzer composed of a standard CCD-video camera, a

- 15 video digitiser and a personal computer with WinGrain software. For every bread, the results of pore diameter in mm and pore homogeneity were calculated as an average of measurements from 10 slices of bread. The pore homogeneity was expressed in % of pores that are larger than 0.5 times the
- 20 average of pore diameter and smaller than 2 times the average diameter.

### (iv) Determination of firmness

25 The firmness of bread, expressed as  $N/dm^2$ , was measured by means of an Instron UTM model 4301 connected to a personal computer. The conditions for measurement of bread firmness were:

Load Cell Max. 100 N

30 Piston diameter 50 mm

Cross head speed 200 mm/min

Compression 25% Thickness of bread slice 11 mm

35 The result was an average of measurements on 10 bread slices for every bread.

### (v) Determination of gluten index

Gluten index was measured by means of a Glutomatic 2200 from Perten Instruments (Sweden). Immediately after proofing, 15 g of dough was scaled and placed in the Glutomatic and washed with 500 ml 2% NaCl solution for 10 min. The washed dough was transferred to a Gluten Index Centrifuge 2015 and the two gluten fractions were scaled and the gluten index calculated according to the following equation:

10

Gluten index = (weight of gluten remaining on the sieve x 100)/total weight of gluten

### (vi) Extraction of lipids from dough

15

30 g of fully proofed dough was immediately frozen and freeze-dried. The freeze-dried dough was milled in a coffee mill and passed through a 235 µm screen. 4 g freeze-dried dough was scaled in a 50 ml centrifuge tube with screw lid 20 and 20 ml water saturated n-butanol (WSB) was added. The centrifuge tube was placed in a water bath at a temperature of 100°C for 10 min. after which the tubes were placed in a Rotamix and turned at 45 rpm for 20 min. at ambient temperature. The tubes were again placed in the water bath for 10 min. and turned on the Rotamix for another 30 min. at ambient temperature.

The tubes were centrifuged at 10,000 x g for 5 min. 10 ml of the supernatant was pipetted into a vial and evaporated to 30 dryness under nitrogen cover. This sample was used for HPLC analysis.

A similar sample was fractionated on a Bond Elut Si (Varian 1211-3036). The non-polar fraction was eluted with 10 ml 35 cyclohexan:isopropanol:acetic acid (55:45:1) and evaporated to dryness. This sample was used for GLC analysis.

### (vii) HPLC analysis

Column: LiChrospher 100 DIOL 5  $\mu m$  (Merck art. 16152) 250x4 mm with a water jacket of a temperature of 50°C.

5

Mobile phases:

A: heptan:isopropanol:n-

butanol:tetrahydrofuran:isooctan:water (64.5:17.5:7:5:5:1)

10 B: isopropanol:n-butanol:tetrahydrofuran:isooctan:water (73:7:5:5:10)

The mobile phases contained 1 mmol trifluoroacetic acid per 1 mobile phase and were adjusted to pH 6.6 with ammonia.

15

Pump: Waters 510 equipped with a gradient controller.

#### Gradient:

	Flow	(ml/min)	Time (min) A	(%) B (%)
20	1.0	0	100	0
	1.0	25	0	100
	1.0	30	0	100
	1.0	35	100	0
	1.0	40	100	0

25

Detector: CUNOW DDL21 (evaporative light-scattering); temperature 100°C; voltage: 600 volt; air flow: 6.0 l/min.

Injector: Hewlett Packard 1050; injection volume: 50 µl.

30

The samples for analysis were dissolved in 5 ml chloroform:methanol (75:25), sonicated for 10 min and filtered through a 0.45  $\mu m$  filter.

### 35 (viii) GLC analysis

Perkin Elmer 8420 Capillary Gas Chromatograph equipped with WCOT fused silica column 12.5 m x 0.25 mm coated with 0.1  $\mu m$ 

stationary phase of 5% phenyl-methyl-silicone (CP Sil 8 CB from Crompack).

Carrier: Helium

Injection: 1.5 µl with split

5 Detector: FID 385°C

Oven program:	1	2	3	4
Oven temperature, °C	80	200	240	360
Isothermal time, min	2	0	0	10
Temperature rate, °C/min	20	10	12	

10

Sample preparation: 50 mg non-polar fraction of wheat lipids was dissolved in 12 ml heptane:pyridine (2:1) containing 2 mg/ml heptadecane as internal standard. 500 µl of the solution was transferred to a crimp vial and 100 µl N-methyl-N-trimethylsilyl-trifluoracetamide was added. The mixture was allowed to react for 15 min at 90°C.

Calculation: Response factors for mono-, di- and triglycerides and free fatty acids were determined from 20 reference mixtures of these components. Based on these response factors, the glycerides and the free fatty acids were calculated in wheat lipids.

### 8.2. Baking experiments with lipase 3 in Danish toast bread

25

The effect of adding lipase 3 to a dough for making Danish toast bread was evaluated. The enzyme was added as a freeze-dried preparation on maltodextrin together with the other ingredients. The results of the baking tests are shown in 30 Tables 8.1 to 8.4.

Table 8.1

Lipase LUS/kg flour	0	5,000	15,000	25,000
Specific volume of bread	4.43	4.43	4.22	4.37
Firmness Day 1	35	33	32	30
Firmness Day 7	90	90	85	73

Table 8.2

Lipase LUS/kg flour	0	5,000	15,000	25,000
Average diameter of the crumb pore, mm	2.96	2.33	2.47	2.65
Homogeneity of crumb pore, %	64.9	73.8	66.0	67.1
Porosity, %	85.9	84.7	85.5	85.1
Gluten index, %	42	45.5	55	65

### 5 <u>Table 8.3</u>

Lipase LUS/kg flour	0	5,000	15,000	25,000
Fatty acids, %	0.090	0.148	0.218	0.241
Monoglycerides, %	0.017	0.031	0.035	0.039
Diglycerides, %	0.020	0.036	0.040	0.045
Triglycerides, %	0.790	0.714	0.673	0.622

Table 8.4

Lipase LUS/kg flour	0	5,000	15,000	25,000
Monogalactosyl Diglyceride, %	0.073	0.040	0.025	0.018
Digalactosyl Diglyceride, %	0.244	0.220	0.182	0.127
Digalactosyl Monoglyceride, %	0.008	0.022	0.044	0.054
Phosphatidyl choline, %	0.064	0.073	0.055	0.041
Lysophosphatidyl choline, %	0.164	0.182	0.171	0.165

By the addition of up to about 5,000 LUS/kg flour of the lipase no change in bread volume was observed, but at a 5 higher dosage of lipase 3 there was a tendency to a small but not statistically significant decrease in volume (Table 8.1).

From the results in Table 8.2 it appears that lipase 3 improved the bread crumb homogeneity and that the average 10 diameter of the crumb pores was reduced significantly. The gluten index also clearly correlated to the addition of lipase 3 as an indication of a more firm gluten caused by the modification of the wheat lipid components causing better dough stability and a more homogeneous bread pore structure.

- 15 However, these modifications appeared to be optimal at the addition of 5,000 LUS/kg flour of lipase 3 whereas a higher dosage resulted in a too strong modification of the wheat gluten.
- 20 The results of the GLC and HPLC analyses (Table 8.3) clearly demonstrated that the triglycerides in the dough were hydrolysed. But more interestingly, there was also observed a modification of the glycolipids, monogalactosyl diglyceride

and digalactosyl diglyceride. These components were converted to the more polar components monogalactosyl monoglyceride and digalactosyl monoglyceride. As digalactosyl monoglyceride is a more surface active component than digalactosyl diglyceride it is assumed that this component contributed to the observed improved crumb cell structure and homogeneity. It also appeared that phospholipids like phosphatidyl choline were only modified to a very small extent.

### 10 8.3. Baking experiments with lipase 3 in Danish rolls

The effect of adding lipase 3 to a dough for making Danish rolls was evaluated. The enzyme was added as a freeze-dried preparation on maltodextrin together with the other ingre15 dients. The results of the baking tests are shown in Tables 8.5 to 8.7.

Table 8.5

Lipase 3 LUS/kg flour	0	10,000	20,000	30,000
Specific volume of bread (45 min fermentation)	6.86	7.04	6.35	6.36
Specific volume of bread (65 min fermentation)	8.30	8.59	8.23	8.04
Subjective evaluation of crumb homogeneity	3	5	4	4

Table 8.6

Lipase 3 LUS/kg flour	0	10,000	20,000	30,000
Free fatty acids, %	0.060	0.126	0.173	0.211
Monoglycerides, %	0.028	0.050	0.054	0.063
Diglycerides, %	0.103	0.095	0.110	0.104
Triglycerides, %	0.705	0.561	0.472	0.436

Table 8.7

5

		4		
Lipase 3 LUS/kg flour	0	5,000	15,000	25,000
Digalactosyl Diglyceride, %	0.204	0.187	0.154	0.110
Digalactosyl Monoglyceride, %	0.007	0.026	0.047	0.074
Phosphatidyl choline, %	0.077	0.078	0.077	0.063
Lysophosphatidyl choline, %	0.153	0.161	0.162	0.150

It is apparent from the results shown in Table 8.5 that the addition of lipase 3 does not significantly increase the volume of the rolls. Furthermore, lipase 3 was found to 10 improve the homogeneity of the crumb.

The GLC and HPLC analyses of the wheat lipids, as shown in Tables 8.6 and 8.7, demonstrated the modification of these lipids.

5.4

### EXAMPLE 9

### Dough improving effect of glycerol oxidase and lipase

- 5 The effect of glycerol oxidase and lipase (separately or in combination) on dough strength was studied in a dough prepared according to the AACC Method 54-10. The dough was subjected to extensiograph measurements (Barbender Extensiograph EXEK/6) also according to AACC Method 54-10 with and 10 with out the addition of glycerol oxidase from Aspergillus japonicus combined with lipase from Aspergillus oryzae (GRINDAMYLTM EXEL 16, Bakery Enzyme, Danisco Ingredients). The dough with out addition of enzymes served as a control.
- 15 The principle of the above method is that the dough after forming is subjected to a load-extension test after resting at 30°C for 45, 90 and 135 minutes, respectively, using an extensigraph capable of recording a load-extension curve (extensigram) which is an indication of the doughs resistance 20 to physical deformation when stretched. From this curve, the
- resistance to extension, B (height of curve) and the extensibility, C (total length of curve) can be calculated.

  The B/C ratio (D) is an indication of the baking strength of the flour dough. The results of the experiment are summarized
- 25 in Table 9.1 below.

30

Table 9.1: Extensigraph measurements of dough supplemented with glycerol oxidase and lipase

Sample (per kg flour)	Resting time (min)	B-value	C-value	D=B/C
Control	45	220	192	1.15
500 LUS lipase	45	225	190	1.18
1000 U glycerol oxidase	45	300	195	1.54
500 LUS lipase + 1000 U Glycerol oxidase	45	350	198	1.77
Control	90	240	196	1.22
500 LUS lipase	90	245	195	1.16
1000 U Glycerol oxidase	90	330	190	1.74
500 LUS lipase + 1000 U Glycerol oxidase	90	380	192	1.98
Control	135	260	188	1.38
500 LUS lipase	135	265	190	1.39
1000 U Glycerol oxidase	135	380	188	2.02
500 LUS lipase + 1000 U Glycerol oxidase	135	410	190	2.15

<sup>5</sup> When the results from the above experiments are compared with regard to the differences between the control dough and the glycerol oxidase supplemented dough it appears that glycerol oxidase clearly has a strengthening effect. The B/C ratio was increased by 34%, 43% and 46% after 45, 90 and 135 minutes of 10 resting time respectively.

The addition of lipase only did not have any effect on the B/C ratio.

However, when supplementing the dough with a combination of glycerol oxidase and lipase, a further increase in the B/C ratio was seen as compared to bread prepared from dough supplemented with glycerol oxidase only. The B/C ratio was increased by 54%, 62% and 56% after 45, 90 and 135 minutes respectively. This clearly indicates that the combined use of these two enzymes in the preparation of bread products has an enhancing effect on the baking strength.

10

### EXAMPLE 10

## Improvement of the specific volume of bread prepared from dough supplemented with glycerol oxidase and lipase

15

The effect of using glycerol oxidase and lipase (separately or in combination) on the specific bread volume and the crumb homogeneity was tested in a baking procedure for Danish rolls with a dough prepared as described in example 8. Glycerol

- 20 oxidase from Aspergillus japonicus and lipase 3 from Aspergillus tubigensis was added to the dough in different amounts. Dough without the addition of enzymes served as control. The fully proofed dough was baked at 220°C for 18 minutes with 12 seconds steam in a Bago-oven. After cooling
- 25 the rolls were weighed and the volume of the rolls were measured by the rape seed displacement method. The specific bread volume was determined as the volume of the bread (ml) divided by the weight of the bread (g). The crumb homogeneity was evaluated subjectively on a scale from 1 to 7, where 1 =
- 30 course inhomogeneous and 7 = nice homogeneous.

  The results from this experiment are summarized in Table 10.1 below.

35

Table 10.1: Specific volume and crumb homogeneity in bread supplemented with lipase and glycerol radius.

Sample (per kg flour)	Specific vo- lume (ml/g)	Crumb homo- geneity
Control	5.45	1
1,000 U glycerol oxidase	6.75	2
10,000 LUS lipase	5.65	4
10,000 LUS lipase + 1,000 U glycerol oxidase	7.25	7

5

As can be seen in the above Table 10.1, the use of glycerol oxidase in the preparing of bread, significantly increased the bread volume (24%) as compared to bread prepared from a similar dough not supplemented with this enzyme. Addition of glycerol oxidase did not improve the crumb homogeneity significantly.

The use of lipase in the preparing of bread did not increase the specific volume of the bread, however a highly increased 15 pore homogeneity was observed.

The combined use of glycerol oxidase and lipase increased the specific volume of the bread with 33% as compared to bread prepared from a similar dough not supplemented with any of 20 the two enzymes.

In addition, the crumb homogeneity was highly improved by the combined use of lipase and glycerol oxidase as compared to the control bread and the breads prepared from dough supple25 mented with lipase and glycerol oxidase respectively.

This clearly indicates that the combination of lipase and glycerol oxidase in the preparation of bread has a synergis-

tic effect and significantly enhances the shape and appearance of the finished bread product.

### EXAMPLE 11

5

## Hydrolysis of triglycerides and formation of glycerol in dough supplemented with lipase

In order to study the hydrolysis of triglycerides and the formation of glycerol in a proofed dough supplemented with lipase, a dough for Danish rolls was prepared in the same manner as described in example 8. Different amounts of lipase (GRINDAMYLTM EXEL 16) was added to the dough, and the total lipid from the fully proofed dough was extracted and analyzed by gas chromatography as described above.

Table 11.1: Triglycerides and glycerol in a dough as a function of lipase addition

Lipase addition (GRINDAMYLTM EXEL 16) (LUS per kg flour)	Glycerol	Triglycerides
0	2.2	7.88
500	2.2	6.22
1,250	2.4	5.99
2,500	2.8	5.37
3,750	2.9	5.47
5,000	3.0	5.55
7,500	3.1	5.03
10,000	3.0	4.39

20

From the above experiment it is clear that the addition of

lipase to a dough has a hydrolyzing effect on the triglycerides present in the dough, which is seen as a decrease in the triglyceride content as function of the increased lipase addition. The resulting level of glycerol increases as a function of the lipase addition.

These results suggests, that the improvement of the B/C ratio and the specific bread volume in bread prepared from dough supplemented with both glycerol oxidase and lipase, as was seen in example 9 and 10, could be due to that lipase addi-

10 tion to a dough is generating glycerol which further can act as substrate for glycerol oxidase.

60 SUMMARY PARAGRAPHS The present invention is defined in the claims and the accompanying description. 5 For convenience other aspects of the present invention are presented herein by way of numbered paragraphs. 1. A method of improving the rheological properties of a 10 flour dough and the quality of the finished product made from the dough, comprising adding to the dough 10 to 10,000 units of a glycerol oxidase per kg of flour. 2. A method according to paragraph 1 wherein the glycerol 15 oxidase is derived from an organism selected from the group consisting of a bacterial species, a fungal species, a yeast species, an animal cell and a plant cell. 3. A method according to paragraph 2 wherein the fungal 20 species is selected from the group consisting of an Aspergillus species, a Neurospora species and a Penicillium species. 4. A method according to paragraph 1 wherein the resistance 25 to extension of the dough in terms of the ratio between resistance to extension (height of curve, B) and the extensibility (length of curve, C), i.e. the B/C ratio, as measured by the AACC method 54-10 is increased by at least 10% relative to that of an otherwise similar dough not 30 containing glycerol oxidase. 5. A method according to paragraph 1 wherein the finished product is selected from the group consisting of a bread product, a noodle product and an alimentary paste product. 35 6. A method according to paragraph 1 where at least one further enzyme is added to the dough ingredients, dough additives or the dough.

62 further enzyme is added to the dough ingredients, dough additives or the dough. 15. A method according to paragraph 14 wherein the further 5 enzyme is selected from the group consisting of a cellulase, a hemicellulase, a starch degrading enzyme, an oxidoreductase, and a protease. 16. A method according to paragraph 8 wherein the amount of 10 lipase is in the range of 10 to 100,000 LUS per kg of flour. 17. A method according to paragraph 8 wherein the lipase is derived from an organism selected from the group consisting of a bacterial species, a fungal species, a yeast species, an 15 animal cell and a plant cell. 18. A method according to paragraph 17 wherein the lipase is derived from an Aspergillus species. 20 19. A method according to paragraph 18 wherein the Aspergillus species is selected from the group consisting of A. tubigensis, A. oryzae and A. niger. 20. A method according to paragraph 8 wherein at least 10% of 25 the galactosyl diglycerides normally present in a flour dough is hydrolysed to the corresponding galactosyl monoglycerides. 21. A dough improving composition comprising a glycerol oxidase and at least one further dough ingredient or dough 30 additive. 22. A composition according to paragraph 21 wherein the further dough additive is selected from the group consisting of a substrate for glycerol oxidase and a lipase. 35 23. A composition according to paragraph 22 which is a premixture useful for preparing a baked product or in making a noodle product or an alimentary paste product.

63 24. A composition according to paragraph 21 which comprises an additive selected from the group consisting of an emulsifying agent and a hydrocolloid. 5 25. A composition according to paragraph 24 wherein the hydrocolloid is selected from the group consisting of an alginate, a carrageenan, a pectin and a vegetable gum. 10 26. A composition according to paragraph 21 wherein the amount of glycerol oxidase is in the range of 10 to 10,000 units per kg flour. 27. A composition according to paragraph 21 or 26, comprising 15 as the further dough additive a lipase in an amount which is in the range of 10 to 100,000 LUS per kg flour. 28. Use of a glycerol oxidase for improving the rheological properties of a flour dough and the quality of the finished 20 product made from the dough. 29. Use according to paragraph 28 wherein the improvement of the rheological properties include that the resistance to extension of the dough in terms of the ratio between 25 resistance to extension (height of curve, B) and the extensibility (length of curve, C), i.e. the B/C ratio, as measured by the AACC method 54-10 is increased by at least 10% relative to that of an otherwise similar dough not containing glycerol oxidase. 30 30. Use of a glycerol oxidase and a lipase in combination for improving the rheological properties of a flour dough and the quality of the finished product made from the dough. 35 31. Use according to paragraph 30 wherein the improvement of the rheological properties of the dough include that the resistance to extension of the dough in terms of the ratio between resistance to extension (height of curve, B) and the

64 extensibility (length of curve, C), i.e. the B/C ratio, as measured by the AACC method 54-10 is increased by at least 10% relative to that of an otherwise similar dough that does not contain glycerol oxidase. 5 32. Use according to paragraph 30 wherein the improvement of the quality of the finished product made from the dough is that the average pore diameter of the crumb of the bread made from the dough is reduced by at least 10%, relative to a 10 bread which is made from a bread dough without addition of the lipase. 33. Use according to paragraph 30 wherein the improvement of the quality of the finished product made from the dough is 15 that the pore homogeneity of the crumb of the bread made from the dough is increased by at least 5%, relative to a bread which is made from a bread dough without addition of the lipase. 20 34. Use according to paragraph 30 or 31 wherein the improvement of the rheological characteristics of the dough includes that the gluten index in the dough is increased by at least 5%, relative to a dough without addition of a lipase, the gluten index is determined by means of a 25 Glutomatic 2200 apparatus.

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### INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A. The indications made below relate to the microorganism reform on page 37, line 6	erred to in the description										
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet										
Name of depositary institution The National Collections of Industrial and Marine Bacteria Limited (NCIMB)											
Address of depositary institution (including postal code and country)											
23 St. Machar Drive, Aberdeen, Scotland United Kingdom, AB2 1RY											
Date of deposit 24 February 1997	Accession Number NCIMB 40863										
C. ADDITIONAL INDICATIONS (lowe blank if not applicable	(c) This information is continued on an additional sheet										
As regards the respective Patent Offices of the respective designated states, the applicants request that a sample of the deposited microorganisms only be made available to an expert nominated by the requester until the date on which the patent is granted or the date on which the application has been refused or withdrawn or is deemed to be withdrawn.											
D. DESIGNATED STATES FOR WHICH INDICATIO	NS ARE MADE (if the indications are not for all designated States)										
E. SEPARATE FURNISHING OF INDICATIONS (leave	: blank if not applicable)										
The indications listed below will be submitted to the International Number of Deposit')											
For receiving Office use only	For International Bureau use only										
This sheet was received with the international application	This sheet was received by the International Bureau on:										
Authorized officer  Jette Cordes Paulsen	Authorized officer										

### SEQUENCE LISTING

5	(1) GENERAL INFORMATION:
	(i) APPLICANT:
	(A) NAME: DANISCO A/S
	(B) STREET: Langebrogade 1
	(C) CITY: Copenhagen
10	(E) COUNTRY: Denmark
	(F) POSTAL CODE (ZIP): 1001 K
	(G) TELEPHONE: +45 32 66 22 00
	(H) TELEFAX: +45 32 66 21 67
15	(ii) TITLE OF INVENTION: Cloning and use of Lipase 3 gene from Aspergillus tubigensis
	(iii) NUMBER OF SEQUENCES: 9
20	(iv) COMPUTER READABLE FORM:
	(A) MEDIUM TYPE: Floppy disk
	(B) COMPUTER: IBM PC compatible
	(C) OPERATING SYSTEM: PC-DOS/MS-DOS
25	(D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)
	(2) INFORMATION FOR SEQ ID NO: 1:
	(i) SEQUENCE CHARACTERISTICS:
30	(A) LENGTH: 25 amino acids
	(B) TYPE: amino acid
	(C) STRANDEDNESS: single
	(D) TOPOLOGY: linear
35	(ii) MOLECULE TYPE: peptide
	(v) FRAGMENT TYPE: N-terminal
	(vi) ORIGINAL SOURCE:
10	(A) ORGANISM: Aspergillus tubigensis

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1: Ser Val Ser Thr Ser Thr Leu Asp Glu Leu Gln Leu Phe Ala Gln Trp 5 10 15 5 Ser Ala Ala Tyr Xaa Ser Asn Asn 20 25 (2) INFORMATION FOR SEQ ID NO: 2: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 7 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single 15 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide (v) FRAGMENT TYPE: internal 20 (vi) ORIGINAL SOURCE: (A) ORGANISM: Aspergillus tubigensis 25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2: Val His Thr Gly Phe Trp Lys 5 30 (2) INFORMATION FOR SEQ ID NO: 3: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 14 amino acids 35 (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: peptide 40 (v) FRAGMENT TYPE: internal (vi) ORIGINAL SOURCE: (A) ORGANISM: Aspergillus tubigensis 45

	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:	
5	Ala Trp Glu Ser Ala Ala Asp Glu Leu Thr Ser Lys Ile Lys 1 5 10	
	(2) INFORMATION FOR SEQ ID NO: 4:	
10	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 20 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
15	<pre>(ii) MOLECULE TYPE: other nucleic acid   (A) DESCRIPTION: /desc = "Oligonucleotide"</pre>	
20	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:	
	TTCCARAANC CNGTRTGNAC	20
25	(2) INFORMATION FOR SEQ ID NO: 5:	
30	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 18 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
35	<pre>(ii) MOLECULE TYPE: other nucleic acid   (A) DESCRIPTION: /desc = "Oligonucleotide"</pre>	
40	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:	
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45	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 17 base pairs	

(B) TYPE: nucleic acid

	<pre>(C) STRANDEDNESS: single (D) TOPOLOGY: linear</pre>														
5	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>														
	(10)														
10	A LA COMPAGN PROGRAMMAN AND AND AND AND AND AND AND AND AND A														
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:														
	GCVGCHSWYT CCCAVGC														
15	(2) INFORMATION FOR SEQ ID NO: 7:														
	(i) SEQUENCE CHARACTERISTICS:														
	(A) LENGTH: 317 base pairs														
20	(B) TYPE: nucleic acid (C) STRANDEDNESS: double														
	(D) TOPOLOGY: linear														
	(ii) MOLECULE TYPE: other nucleic acid														
25	(A) DESCRIPTION: /desc = "PCR fragment"														
23															
2.0	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:														
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	TCCAACTTGA CATGCACGGC CAACGCCTGT CCATCAGTCG AGGAGGCCAG TACCACGATG	120													
35	CTGCTGGAGT TCGACCTGTA TGTCACTCAG ATCGCAGACA TAGAGCACAG CTAATTGAAC	180													
	AGGACGAACG ACTTTTGGAG GCACAGCCGG TTTCCTGGCC GCGGACAACA CCAACAAGCG	240													
4 0	GCTCGTGGTC GCCTTCCGGG GAAGCAGCAC GATTGAGAAC TGGATTGCTA ATCYTGACTT	300													
40	CATCCTGGRA GATAACG	317													
	(2) INFORMATION FOR SEQ ID NO: 8:														
45	<ul><li>(i) SEQUENCE CHARACTERISTICS:</li><li>(A) LENGTH: 1045 base pairs</li></ul>														

			·	•		nuc DEDN											
			(	D) T	OPOL	OGY:	lin	ear									
5		(ii	) MO	LECU	LE T	YPE:	DNA	(ge	nomi	c)							
		(vi	) OR	IGIN	AL S	OURC	E:										
			(	A) O	RGAN	ISM:	Asp	ergi	llus	tub:	igen	sis					
10		(ix	) FE	ATUR	E:												
			·	•		KEY: ION:		(1	82,	135.	.300	, 34	76	83,	737.	.1045)	
		(ix	) FE	ATUR	E:												
15						KEY: ION:			tide								
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20						KEY: ION:	•			300	o, 3	47	683,	737	10	42)	
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		(xi	) SE	QUEN	CE D	ESCR:	IPTI	ON:	SEQ :	ID NO	D: 8	:					
25																CTG	48
	Met -27	Phe	Ser -25	Gly	Arg	Phe	Gly	Val -20	Leu	Leu	Thr	Ala	Leu -15	Ala	Ala	Leu	
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30	Gly	Ala -10	Ala	Ala	Pro	Ala	Pro -5	Leu	Ala	Val	Arg						
	CCG	ATGT	GAG A	ATGG	TTGG	AT A	GCAC	rgato	G AA	GGT	GAAT	AG	GT (	GTC '	rcg i	ACT	145
35												:	Ser ' l	Val :	Ser '	Thr	
	TCC	ACG	TTG	GAT	GAG	TTG	CAA	TTG	TTC	GCG	CAA	TGG	TCT	GCC	GCA	GCT	193
		Thr	Leu	Asp	Glu	Leu	Gln	Leu	Phe	Ala		Trp	Ser	Ala	Ala		
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	Tyr	Cys	Ser	Asn	Asn 25	Ile	Asp	Ser	Lys	Asp 30	Ser	Asn	Leu	Thr	Cys 35	Thr	
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	Ala	Asn	Ala	Cys	Pro	Ser	Val	Glu	Glu	Ala	Ser	Thr	Thr	Met	Leu	Leu	

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				40	ı				45					50			
5			GAC Asp	Leu		TGTC	ACT	CAGA	TCGC	AG A	CATA	.GAGC	A CA	GCTA	ATTT		340
10	GAA	CAG					e Gl					y Ph				G GAC a Asp 70	389
10						Leu			GCC Ala								437
15									TTC Phe 95								485
20									ACT Thr								533
25									AAG Lys								581
30									ACC Thr								629
									CTG Leu								677
	GAG Glu		GTGA	GTCC	CTT (	CACAA	AGGT	G A	rggad	GCGAC	C AA:	rcgg	GAAC	AGA	CAGTO	CAA	733
10									ATC Ile								781

CAT ATC ACC AGT CAG GGA TCT GGG GCC AAC TTC CGT GTT ACA CAC TTG His Ile Thr Ser Gln Gly Ser Gly Ala Asn Phe Arg Val Thr His Leu

195

190

45 185

									CCC								877
	Asn	Asp	Ile	Val	Pro	Arg	Val	Pro	Pro	Met	Asp	Phe	Gly	Phe	Ser		
5	200					205					210					215	
													666	л <i>С</i> -Ш	СТС	N.C.C	925
	CCA	AGT	CCG	GAA	TAC	TGG	ATC	ACC	AGT	GGC	AAT	GGA	71-	AGI	U a l	Thr	723
	Pro	Ser	Pro	Glu		Trp	Ile	Thr	Ser		ASI	GIA	Ala	361	230	1111	
4.0					220					225					230		
10	666	TCC	CAT	א ייי כי	CDD	GTC	ΔTC	GAG	GGA	ATC	AAT	TCA	ACG	GCG	GGA	AAT	973
									Gly								
	Ala	261	лэр	235	010				240					245			
15	GCA	GGC	GAA	GCA	ACG	GTG	AGC	GTT	GTG	GCT	CAC	TTG	TGG	TAC	TTT	TTT	1021
	Ala	Gly	Glu	Ala	Thr	Val	Ser	Val	Val	Ala	His	Leu	Trp	Tyr	Phe	Phe	
			250					255					260				
																	1045
		ATT						AAT									1045
20	Ala		Ser	Glu	Cys	Leu		*									
		265					270										
	(2)	INF	ORMA'	TTON	FOR	SEO	ID	NO:	9:								
25	(2)	1111	J14 2 1														
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			(2	A) L	ENGT	H: 2	97 a	mino	aci	ds							
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35	-27	rne	-25	O.T.J	9		2	-20					-15				
55																	
	Gly	Ala	Ala	Ala	Pro	Ala	Pro	Leu	Ala	Val	. Ar	g Se	r Val	L Se:	r Th	r Ser	
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40	Thr	Leu	Asp	Glu	Leu	Gln	Leu	Phe	e Ala			Se:	r Ala	a Al		a Tyr	
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	Cys	Ser	Asn			: Asp	ser	. rys	30 30		. ASI	. ne	<u> </u>	. cy 3		r Ala	
45				25	•				50	•							
43		Δla	Cvs	Pro	. Ser	. Val	Glı	ı Glu	ı Ala	a Sei	r Th	r Th	r Me	t Le	u Le	u Glu	
	LIGIT	, ,,,,	. 0,3														

			40					45			50						
5	Phe	Asp 55		Thr	Asn	Asp	Phe 60	Gly	Gly	Thr	Ala	Gly 65	Phe	Leu	Ala	Ala	
J	Asp 70		. Thr	Asn	Lys	Arg 75	Leu	Val	Val	Ala	Phe 80	Arg	Gly	Ser	Ser	Thr 85	
10	Ile	Glu	Asn	Trp	Ile 90	Ala	Asn	Leu	Asp	Phe 95	Ile	Leu	Glu	Asp	Asn 100	Asp	
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	Phe	Phe	Ala	Ile	Ser	Glu	Cys	Leu	Leu	*							